Center of Excellence for Chemical Hydrogen Storage: LANL Tasks and Collaborations

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Project ID # STP61



Overview

Timeline

Project Start Date: FY05

Project End Date: FY09

New Start

Budget

Total project funding (requested)

\$ 8.7 M DOE share

Funding for FY05: \$ 1.2 M

Barriers Addressed

Cost

Weight and volume

Energy efficiency

System life-cycle assessment

Spent material removal

Regeneration processes

Heat removal

Partners

Chemical Hydrogen Storage
Center of Excellence Partners



Objectives

- Identify, research, develop and validate the best chemical hydrogen storage systems to overcome barriers and meet 2010 DOE goals
- Develop materials, catalysts, catalytic processes and new concepts for hydrogen release and regeneration
- Design, synthesize, and test structures and compositions to control thermochemistry of H₂ release and spent fuel regeneration
- Provide experimental tools to Center partners
- Collaborate with Center partners where appropriate and synergistic
- Contribute to demonstration of 1 kg storage system
- Coordinate research, development and engineering efforts within each Tier and overall Center



Three Tier Structure of Center

Tier I:

- Develop borate-to-borohydride (BO to BH) regeneration alternatives and assess economics and life cycle analysis of borohydride/water to hydrogen
 - Millennium Cell, Rohm and Haas, Penn State, Alabama, PNNL, LANL

Tier II:

- Avoid water and thermodynamic sinks. Alternative boron chemistry approaches include polyhedral boranes (B_xH_y), amine-boranes and BCNP chemistry
 - Penn, Penn State, UCLA, Washington, Northern Arizona, Alabama, Internatix, PNNL, LANL

Tier III:

- Beyond boron:
 - Develop concepts for coupled endo/exothermic reactions, Investigate nanomaterials
 - Use heteroatom substitution for thermodynamic control
 - UC Davis, Alabama, Internatix, PNNL, LANL

Core Capabilities:

- Computation, engineering assessment, complex instrumentation (PNNL, LANL, Rohm and Haas, Alabama)
- IP management (IP Management Committee)
- Safety (PNNL, LANL, Northern Arizona)
- Center coordination, meetings, technical planning (LANL)



4

Approach: Tier 1 B-O → B-H

- Engineering Guided Research
 - Data mining on efficient routes for regeneration of B-H from borates
 - Down-selection of new experimental approaches for regeneration of B-H
 - with US Borax, Millennium Cell, PNNL, Penn State, Rohm and Haas
- Investigation of complexation of borates for reduction chemistry/electrochemistry
 - with PNNL, Rohm and Haas
- Electrochemical studies of borates and complexed borates
 - with Rohm and Haas, Penn State



Approach: Tier 2 Novel Boron Chemistry

- Polyhedral Borane Chemistry
 - Validation of leads and assistance for rapid throughput catalyst development for hydrolysis of polyhedral boranes
 - with Intematix, UCLA, PNNL
 - Electrochemical studies of polyhedral boranes
 - with PSU, UCLA
 - Aminolysis of polyhedralboranes
 - with UCLA, PNNL
- Amine-Borane Chemistry
 - Catalyst development for dehydrogenation of amine-boranes; selectivity, activity, non-precious metals
 - with Penn, Ala, NAU, PNNL
 - Mechanisms of amine-borane dehydrogenation/regeneration
 - with Penn, UW, PNNL



Approach: Tier 3 Innovation Beyond Boron

- Coupled chemical reactions
 - Develop new concepts coupling endothermic and exothermic reactions with high hydrogen storage capacity
- Heteroatom substituted organics
 - Develop molecular systems with appropriate thermochemistry for hydrogen release and regeneration
 - Develop new mechanisms for hydrogen release from organic molecules
 - with Alabama, PNNL
- Nanoscale materials
 - B, BN and doped-B, Si nanoparticles for hydrogen storage
 - with UC Davis
 - Metal-based nanoparticles by gas-solid synthesis
 - with Internatix



Technical Accomplishments/ Progress/Results

- New start in FY05 (1/20/05)
- Collaborative R&D project plans developed for all projects
- Preliminary data mining started for Tier 1
- Promising preliminary results already obtained for Tier 2 and 3



Tier 1: First Year Milestones

- Data mining (B-O to B-H)
 - with ROH, MCEL, USB, PNNL, PSU
- Survey of complex borates
 - with PNNL, USB, ROH
- Selection of analytical and electrochemical methods for BH₄ determination.
 - with ROH, MCEL, USB, PNNL, PSU



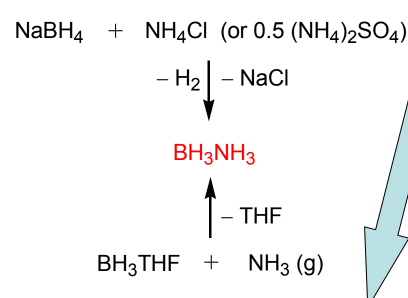
Future Work: Tier 1

- Implement experimental approaches for regenerating B-H, as identified from data mining and downselection
 - with US Borax, Millennium Cell, PNNL, Penn State, Rohm and Haas
- Prepare most-promising borate complexes, as identified from computation and data mining, and determine reduction chemistry
 - with PNNL, Rohm and Haas, Alabama
- Determine electrochemical behavior of borate complexes
- Fabricate electrode materials for electrochemical borate reduction
 - with Rohm and Haas, Penn State



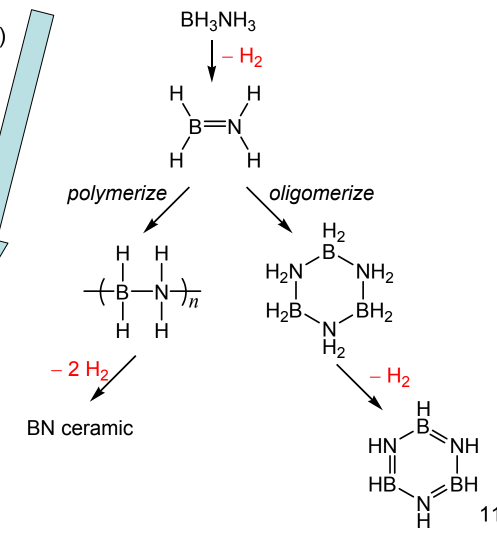
Preliminary Results: Tier 2 Hydrogen Production Using Ammonia-Borane

 Ammonia-borane is easily synthesized from commercially available starting materials. Low molecular weight paired with the loss of several equivalents of hydrogen gives BH₃NH₃ an ideal hydrogen storage capacity of 19.6 wt%.



Polymerization or oligomerization of BH₃NH₃ requires addition of energy (> 80 °C).

Can we find a catalyst to lower the temperature?





Catalyst Development for H₂ Production

$$H$$
 H H_2B H_2B

Manners et al. *JACS* **2003**, *125*, 9424 Fulton, Autrey, et al. *JACS* **2005**, *127*, 3254

Rhodium and other late metal catalysts are expensive

General characteristics of LANL Proprietary catalysts:

Commodity chemicals

Generally air-stable

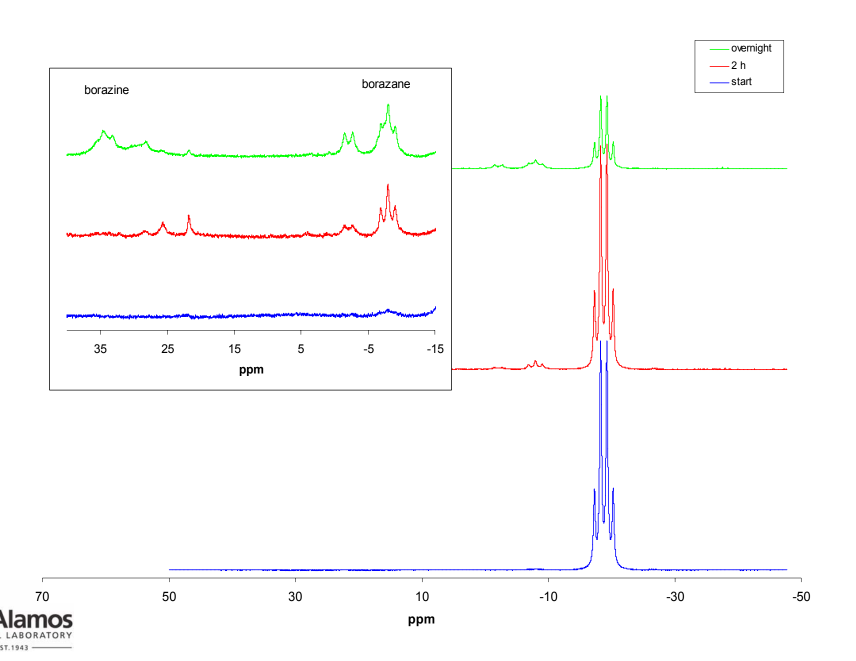
Lower molecular weight than transition metal catalysts

Reactions of BH₃NH₃ with 10 mol% catalyst (or less!) can be observed at room temperature, whereas BH₃NH₃ is stable indefinitely at 25 °C (when rigorously anhydrous, air-free conditions are utilized).

Yes: Our new catalyst system lowers the temperature at which H₂ is evolved from BH₃NH₃

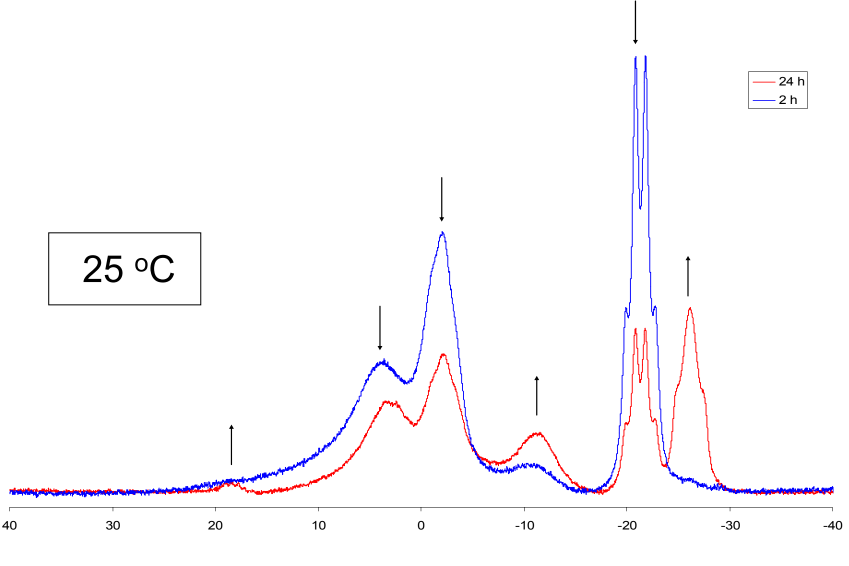


Sample ¹¹B NMR Spectra: Uncatalyzed Thermolysis of BH₃NH₃ at 80 °C



13

Sample ¹¹B NMR Spectra: Catalyzed Dehydrogenation of BH₃NH₃





First Year Milestones

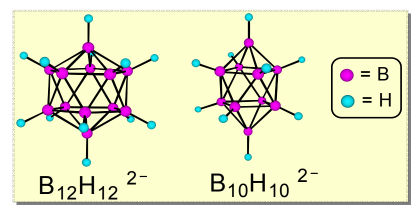
- Determine electrochemical behavior of polyhedral boranes.
 - with PSU, UCLA
- Assist in rapid throughput catalyst development on polyhedral borane hydrolysis/aminolysis
 - with PNNL, Intematix, UCLA
- Optimize reaction conditions for hydrogen evolution from BH₃NH₃
- Use alkyl-substituted, and ²H and ¹⁵N labeled amineboranes to investigate dehydropolymerization mechanism w/ LANL catalysts
- Characterize solid BNH_x products
- Extend dehydrogenation studies to triborane-amine compounds, B,N-substituted analogs, and polyhedral derivatives
 - with/ PNNL, Penn, NAU, UCLA



Polyhedral Boranes

Polyhedral boranes are more stable than borohydride and multiple electron sources

 Optimize catalysts for hydrolysis of polyhedral borane anions using rapid throughput heterogeneous catalyst synthesis and testing



$$Na_2B_{10}H_{10} \cdot 16H_2O \longrightarrow 2 NaBO_2 + 4 B_2O_3 + 21 H_2$$

- Other H2 generation routes
 - Aminolysis
 - Electrochemistry
- New regeneration routes from borate
 - with PNNL, Intematix, PSU, UCLA

Tier 2: Future Work

- Polyhedral borane electrochemistry
 - with PSU, UCLA
- Catalyst development on polyhedral borane hydrolysis, aminolysis
 - with Intematix, PNNL, UCLA
- Mechanisms and catalyst development
- Regeneration of BN systems
 - with PNNL, Penn, NAU, UCLA
- Compare reactivity parameters with computation
 - with PNNL, Alabama
- Determine best practical routes to amine-boranes
 - · with ROH, US Borax, PNNL
- Engage all relevant Center partners to optimize regeneration process and life-cycle for integrated B-N hydrogen storage system(s)
 - Integrated amine-borane hydrogen storage system for engineering and economic assessment.



Preliminary Results: Tier 3

- Coupled reactions to drive hydrogen release
 - Have developed LANL-proprietary concepts for novel means of driving hydrogen-releasing reactions
 - Hypothetical hydrogen capacity > 6 wt%, >0.09 kg
 H₂/liter
 - Modeling, demonstration work begun
- Nanoparticles for hydrogen storage
 - Apparatus developed and demonstrated for preparing nanoparticles by "physical" means (ablation of target, gas-phase selection)
 - Installation of molecular hydrogen doser, atomic hydrogen doser, temperature-programmed-desorption nearing completion



Preliminary Results: Tier 3 Heteroatom-Substituted Organic Compounds

 N-substituents make C-H bond act as "organic hydride"

$$\begin{array}{ccc}
R & H \\
R & R'
\end{array}$$

$$\begin{array}{c|c} R & & & \\ & & & H^{\Theta} \\ R & & & C & R' \\ R & & & R \end{array}$$

- In laboratory test, mixture of "organic hydride" plus acid is unreactive ...
- H + acetic acid

 CH₃

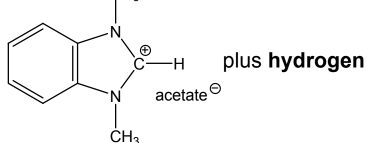
 CH₃

 catalyst

 CH₃

 CH_3

- ... until treated with catalyst
- Path forward: Improve capacity





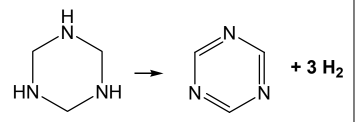


Future Work: Tier 3

- Heteroatom-substituted organic compounds: Demonstrate principle (Y1), useful H₂ release rate (Y2)
 Remove "dead weight" and raise capacity (Y4-5)
- Coupled reactions: Prove hydrogen release (Y1), demonstrate useful H₂ release rate (Y2), using non-precious metals (Y3), pursue to demonstration (Y4-5)
- Nanoparticles: Demonstrate
 working apparatus (Y1), prepare
 nanoparticles rich in B, alloying
 metals, N (Y2); establish ability to
 hydrogenate, dehydrogenate (Y4-5)

Target:

Remove everything except N groups and "hydridic" CH



(material capacity > 6.9 wt% H₂)

U. Alabama has concepts to stabilize, immobilize these structural units



Center Coordination

Objectives

- Real collaboration and information sharing within Tiers
- Collaborative project structure
- Share background information
- Foster joint discovery/inventions
- Reward and manage success

Status

- Joint development projects (Center projects) defined
- Framework for IP management developed and being signed
 - Defines management of joint inventions
 - Enables technology transfer
- Website developed
- Center project meetings, conference calls
- Monthly center-wide "newsletter"



Task	Year 1	Year 2	Year 3	Year 4	Year 5	
1.1 Data Mining		0	ptions ider	tified and o	ownselect	ed
1.1 Complexation of Borate	Laborato	ry studies		Optir	nization	
1.1 Electrochemical Studies				Electrod	e fabrication	ļ φn
2.1 Catalysts for PHB Hydrolysis		Assist wit	h rapid scr	eening Develop	leads	
2.1 Catalysts for PHB Aminolysis			Ass	st with rap	d screenin Develop le	
2.2 PHB Electrochemical Work					2 2 7	
2.3 Catalysts: AB Dehydrogenation				_		
2.4 Regeneration for AB				Ор	timization	
2.4 AB Mechanistic Work						
3.1 Coupled Reactions			Syst	em 1 ➤ Systen	1 2	
3.2 Heteroatom Substituted Organics			Syst	em 1 System	2	
3.3 Nanoscale Materials						
3.4 Main Group Hydrides			—			



Go/No Decision Point



Hydrogen Safety

The most significant hydrogen hazard associated with this project is:

Inadvertant overpressurization of reaction or storage vessels with hydrogen

"Chemical Hydrogen Storage" requires working with chemicals and reactions that evolve hydrogen. In closed vessels, evolved hydrogen may accumulate, resulting in explosion and fire hazards.



Hydrogen Safety

Our approach to deal with this hazard is:

Do not close reaction or storage vessel until certain that the amount of hydrogen that could be evolved can be safely contained by the vessel. If necessary, maintain vent to inert atmosphere. Use relief valves.



Publications and Presentations

- Patent disclosure submitted for new class of amine-borane dehydrogenation catalysts. Provisional patent to be submitted June '05
- Poster presentation, IPHE Hydrogen Storage, Lucca, Italy (June '05)
- Present poster at Inorganic Chemistry Gordon Research Conference (July '05, Newport, RI)
- Two abstracts submitted for American Chemical Society, FUEL Division Hydrogen Storage symposium (August '05, National ACS Meeting, Washington, D.C.)
- Present invited lecture FECHEM 16 (September '05, Budapest, Hungary).
- Paper submitted 3/05 "Hydrogen Evolution from Organic 'Hydrides' "by D. Schwarz et al.
- Provisional patents filed on heteroatom-substituted organic compounds

